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- (54) Negative-working image recording material
- (57) A regative-working image recording material disclosed, which comprises a substance which about slight to generate heat, a water-insoluble and aquoous alkali-soluble resh, and aphenol derivative having in 8 molecule from 4 to 8 benzers nucle, at least one phenolic hydroxyl group and at least two groups represented by the following general formula (f):

wherein R<sup>1</sup> represents a hydrogen atom, an alkyl group or an acyl group. The negative-working image recording material can perform recording independent of the emission wavelength of the exposing light source, particularly with light in the range of from near infrared to infrared (heat radiation).

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#### Description

#### FIELD OF THE INVENTION

The present invention relates to a negative-working image recording material which can be used as an offset printing master. More particularly, the present invention relates to a lithographic printing plate for so called direct plate making process comprising directly making plate from digital signal such as computer signal.

#### BACKGROUND OF THE INVENTION

As systems for directly making plate from digital data from computer there have been heretofore proposed (1) an electrophotographic process, (2) a photocolymentzation process which comprises exposure to Ar laser and post-processing in combination, (3) a process employing a lainnate of a photosensitive resin and a silver sail photographic material, (4) a process employing a silver master, (5) a process which comprises destruction of a silicone rubber layer by discharge or laser sito.

In the bregoing electrophotographic process (1), the processing required for charping, exposure and development is complicated, and the apparatus required is complicated, entire, therither, the recoping photophymicration process (2) requires a post-heating step. This process also requires a high sensitivity plate-making material which is difficult to handle in dwighful to stippin toom).

Furthermore, the foregoing processes (3) and (4) each employ a silver salt that requires a complicated processing adding to cost. Further, the foregoing process (5) can attain a relatively high perfection but leaves something to be desired in the removal of sillicone residual left on the eurface of the printing plate.

On the other hand, laser has made remarkable progress in recent years. In particular, solid laser and semiconductor laser which emit light in the range of near intraced to infrared are now easily available; with a high output and a small size. These lasers are very useful as exposure light sources for use in direct plate making from digital data from computer, etc. However, most practically useful photosersities image recording materials absorb light in the wavelength range of not higher than 450 nm and thus cannot be exposed to these lasers for image recording. Thus, an image recording materials winkt can perform recording independent of the wavelength of the cooper light.

As a technique enabling recording independent of the wavelength of the exposure fight there is disclosed a positivesowinking image recording material comprising a compound which decomposes when acted only light or heat (a.g., diszonium compound), a particulate substance which can absorb light and convert it to heat and a binder in JP-A-S2-113219 (The Ism "JP-A" as used herein meens an "unexamined published Japanese patient application"). When heated, this positive-working image recording material undergoes decomposition of a diszonium compound to record an image.

Further, JP-A-58-148792 discloses a positive-working photosensitive heat-sensitive recording material comprising as sesential components a particulate thermoplastic resin, a photo-heat convertion substance and a photo-orosalintative substance, leg., diszorium compound). This type of a recording resterial fullizes a phenomenon that is particulate thermoplastic resin forms an image when acted on by heat and a photo-crosslintatic substance undergoes direct decomposition when acted on by light to give an image with an enhanced durability.

The direct plate-making process comprises scarning with a beam from a laser source to write an image. In this process, a negative-working material is preferably used because if can chorant he writing first. Neweyer, no regative-working material having good recording properties has herefolize been known which can perfect material having good recording properties has herefolize been known which can perfect material having good recording properties has herefolize been known which can perfect material having an emission wavelength range of from next intraved to infrared as an exposing fight source for use in the direct plate-making from digital data from computers of the properties of the list of the properties of the prope

# SUMMARY OF THE INVENTION

It is therefore an object of the present invention to provide a negative-working image recording material which can so perform recording independent of the emission wavelength of the exposing light power, particularly a negative-working image recording material which can perform recording with light in the wavelength range of from near infrared to infrared (heat radiation).

it is another chipact of the present invention to provide a negative-working lithographic printing plate for heat most writing type direct plate making which can perform recording by means of a sold laser or semiconductor laser (heat as mode) having an emission wavelength range of from near infrared to infrared to make a plate directly from digital data from commutator or the like and make the direct accolarision of conventional processors or printers.

The foregoing objects of the present invention can be accomplished by the following constitution (1) of the present invention:

#### (1) A negative-working image recording material comprising:

- (a) a substance which absorbs light to generate heat;
- (b) a water-insoluble and aqueous alkali-soluble resin; and
- (c) a phenol derivative having in its molecule from 4 to 8 benzene nuclei, at least one phenolic hydroxyl group and at least two groups represented by the following general formula (i):

### wherein R1 represents a hydrogen atom, an alkyl group or an acyl group.

Further, the present invention has the following preferred embodiments (2) to (12):

(2) A negative-working image recording meterial comprising a substance which absorbs light to generate heat, a water-insoluble and aqueous a sikali-actuble resin and a phenot derivative traving in its molecule from 4 be benzere nuclei, at least one phenotic hydroxyl group and at least two groups represented by the general formula (I), characterized in that said vater-insoluble and aqueous alkali-soluble resin is a resin having a phenotic hydroxyl group or cleficiality insenturated point.

- (3) A negative-working image recording material comprising a support having thereon a substance which absorbs infrared rays or near infrared rays to generate heat, a water-insoluble and aqueous alkali-soluble ratio having a phenolic hydroxy group or olderlinasity unsaturated bond, and a phenol derivative having in its molecula from 4 to 8 benzane ruclei, at least one phenolic hydroxyl group and at least two groups represented by the general formula (b) shown above.
- (4) The image recording material according to the embodiment (3), wherein as said substance which absorbs infrared rays or near infrared rays to generate heat there is used a dve:
- (5) The Image recording material according to the embodiment (3), wherein as said substance which absorbs infra-
- red rays or near infrared rays to generate heat there is used a pigment;

  (6) The image recording material according to the embodiment (3), wherein as said substance which absorbs infra-
- red rays or near infrared rays to generate heat there is used carbon black;
- (7) The image recording material according to the embodiment (3), wherein as said water-insoluble and aqueous alkal-soluble resin having a phenotic hydroxyl group or olefinically unsaturated bond there is used a novoluk resin: (3) The image recording material according to the embodiment (3), wherein as said water-insoluble and aqueous alkal-soluble resin having a phenotic hydroxyl group or olefinically unsaturated bond there is used an acrylic resin having an alkid or uso:
- (9) The Image recording material according to the embodiment (3), wherein as said water-insoluble and aqueous alkali-soluble resin having a phenolic hydroxyl group or olefinically unsaturated bond there is used an acrylic resin
  - having a phenolic hydroxyl group; (10) The Image recording material according to the embodiment (3), wherein as said support there is used a polyrester film:
- (11) The image recording material according to the embodiment (3), wherein as said support there is used an aluminum plate:
  - (12) A negative-working image recording process, which comprises exposing a negative-working image recording material according to the embodiment (3) to Infared or near Infared rays emitted by a laser, and then developing the negative-working image recording material with an aqueous alkali;
- (13) The negative-working image recording process according to the embodiment (12), wherein R<sup>1</sup> in the general formula (I) defined in the embodiment (3) is an alkyl group;
  - (14) The negative-working image recording process according to the embodiment (12), wherein R<sup>1</sup> in the general formula (f) defined in the embodiment (3) is a methyl group;
  - (15) A negative-working image recording process, which comprises exposing a negative-working image recording material according to the embodiment (3) to infrared rays or near infrared rays entired to a less; subjecting the negative-working image recording material to heat treatment, and then developing the negative-working image recording material with an accessor shall.

## DETAILED DESCRIPTION OF THE INVENTION

The inventors made extensive studies. As a result, it was found that the combined use of a phenol derivative having undecude from 4 to 8 benzene raudei, at least one phenolic hydroxyl group and at least two groups represented by the general formula (I) with a water-insoluble and appeaus alleai-soluble reals and a substance which steaders bight to generate heat can provide a negative-working image recording material which can perform recording independent of the wavelength of the exposing light source, particularly a negative-working image recording material which can perform recording integers and recording material which can perform recording material which rec

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form recording with light in the range of from near infrared to infrared. Thus, the present invention has been worked out. The present invention comprises converting energy of exposing light to heat energy which is then used to cause condensation reaction by a phenol derivative, thereby recording an image.

The present invention is advantageous in that the use of a phenol derivative having in its molecular from 4 to 8 benunction provides a high sensitivity and a high first strengton on the exposed area as compared with the use of a phenol derivative having in its molecule only from 1 to 3 benzene rucide. If a phenol derivative having in its molecule not less than 9 benzene ruciel is used, it is disadvantageous in that the developability is deteriorated and the resulting insea can be selsify stained.

In the present invention, if the phenol derivative has two or more phenolic hydroxyl groups, it is advantageous in the developability after exposure. If the phenol derivative has bur or more groups represented by the general formula (i), it is advantageous in that the film strength on the exposal area is raised.

In the present invention, if the substance which absorbs light to generate heat is a substance which absorbs infrared rays or near infrared rays to generate heat, it is advantageous in that the heat mode recording can be performed fairly.

In the present invention, if the substance which absorbs inflared rays or near inflared rays to generate is a dys. it is advantageous in the developability after exposure. Further, if the substance which absorbs infrared rays or near inflared rays to generate heat is a pigment, it is advantageous in that the resulting sensitivity is good. Further, if the substance which absorbs infrared rays or near inflared rays to generate heat is carbon black, it is advantageous in that the absorption weredength range is wide and the resulting sometivity is limited.

In the present invention, as the support there can be advantageously used a polyester film to give a lighter weight, wher, an aluminum plate can be advantageously used to give a better dimensional stability.

The negative-working image recording material of the present invention is preferably applied to a negative-working image recording process which comprises supposing the negative-working image recording material to infrared rays or near infrared rays emitted by a laser, and then developing the negative-working image recording material with an aque-

Preferred examples of the negative-working image recording material of the present invention include a negativeworking heat-sensitive lithographic printing plate and a negative-working lithographic printing plate for heat mode writing two direct olate makind.

The heat mode writing is a mode of writing in which a proper heat radiation source is controlled according to digital data to perform recording on the image recording material. As the heat radiation source there may be used a thermal head for use in fractional, e.t. or a Laser emitting intraved rays or near infrared rays. If the thermal head is used, the resulting image has a low resolving power. Therefore, as the heat radiation source for direct plate making there is preferably used a laser emitting infrared rays or near infrared rays.

The present invention comprises converting light to heat via a photo-heat conversion substance, which heat is then used to cause crosslinking recipion. Thus, the present invention is essentially a photoensitive recording material. Preferred examples of radiations employable in the present invention include ultraviolet radiation, visible radiation, and infrared radiation. Among these radiations, infrared radiation is commonly called heat radiation. Thus, if recording is effected on the recording material of the present invention by infrared radiation to the recording material of the present invention may be referred to a heat-exercisive recording material.

The present invention will be further described hereinafter.

Preferred examples of the alfyl group represented by R<sup>2</sup> in the foregoing general formula (f) include Ct<sub>1-4</sub> alfyl group, ps.th as methyl group, a tryl group, ps.th group such as methyl group, sechtyl group, ps.th group, include group, sechtyl group, and thurly group. Preferred examples of the acyl group represented by R<sup>2</sup> in the foregoing general formula (f) include formly group, acyl group, sechly group, burly group, and they group, and they group, ps.th as Ct<sub>1-4</sub> substituted as alley group such as methoxyethyl group, methoxypropyl group, hydroxyethyl group and hydroxypropyl group may be used.

The phenol derivative employable in the present invention can be obtained by reacting a known phenol compound such as those described in JP-A-1-28994S, JP-A-3-17985S, JP-A-3-20265K, JP-A-3-17985S, JP-A-3-20265K, JP-A-3-1985S, JP-A-3-2026K, JP-A-3-1982SS, and JP-A-5-2244G9 with formaldatyde in a strongly alkaline medium at a temperature of from about 0°C to 80°C, pretenably from 10°C to 80°C to 1 to 30 hours. Thus, a phenol derivative wherein R<sup>1</sup> is a hydrogen about an obtained. Thereafter, the phenol derivative where of R<sup>2</sup> is a hydrogen about an obtained may be receded with a 2, alcohol, substituted adorbl, add halde or acid enhydride under acidic conditions at a temperature of from 0°C to 80°C for 1 to 30 hours to obtain a phenol compound wherein R<sup>1</sup> is an algly or acyd. The emperature which the phenol derivative is reacted with an alcohal condition of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition or add of the phenol derivative is reacted with an alcohal condition of the phenol derivative is reacted with an alcohal condition of the phenol deriva

Examples of the phenoi derivative to be used in the present invention include compounds represented by the tich lowing general formulae (II) to (I)b, but he present invention should not be construed as being limited thereo. These phenol derivatives may be used singly or in admixture. The amount of such a phenol derivative to be used is from 0.2 to 60% by welful, preferably from 0.5 to 20% by weight based on the veight of the photosensitive compositions.

A compound having from 1 to 3 benzene nuclei, a phenolic hydroxyl group and a group represented by the general formula (i) causes the drop of inking propriy and development latitude. It is thus desirable that the photosensitive composition of the present invention be substantially free of such a compound. More preferably, the content of such a compound is not more than 5% by weight, even more preferably not more than 3% by weight, and most preferably 0% by weight.

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$$(Y)_{b} \qquad (OH)_{m} \qquad (R^{3})_{q}$$

$$(Y)_{c} \qquad (OH)_{n} \qquad (Y)_{c} \qquad (OH)_{n} \qquad (P^{2})_{p} \qquad (OH)_{n} \qquad (OH)_{$$

$$(R^{2})_{q} \xrightarrow{(OH)_{1}} (R^{10})_{u} \xrightarrow{(OH)_{1}} (R^{2})_{p} \xrightarrow{(Y)_{Y}} (Y)_{x} \xrightarrow{(OH)_{m}} (R^{2})_{q} \xrightarrow{(OH)_{n}} (R^{10})_{t}$$

$$(Y)_{a} \qquad (R^{2})_{p} \qquad (DH)_{1}$$

$$(QH)_{a} \qquad (R^{3})_{q} \qquad (R^{3})_{s} \qquad (QH)_{h} \qquad ($$

$$\begin{array}{c} (Y)_{B} \\ (R^{2})_{p} \\ (R^{2})_{p} \\ (R^{3})_{q} \\ (R^{4})_{r} \\ (R^{5})_{q} \\$$

o (VIII)

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wherein R<sup>2</sup> to R<sup>4</sup>, R<sup>17</sup> and R<sup>18</sup> each represents a hydrogen atom, a halopen atom, an alkyl group or an alkoy group, R<sup>8</sup> and R<sup>13</sup> to R<sup>18</sup> each represents a hydrogen atom or an alkyl group, R<sup>18</sup> to R<sup>8</sup> each represents a hydrogen atom or an alkyl group, R<sup>18</sup> to R<sup>8</sup> each represents a raingle bond, an alkyleven, alkenylevne, nephthylene, cathonyl, either or thioether group which may have a substituent, an amide bond or a combination of two or more of these bonds and groups; Y represents a group represented by the general formula (f); a, b, c, d, x and y sech represents an integer of from 0 to 3, with the proviso that the sum of a, b, c, d, x and y is an integer of the group of the general formula (f); a, b, c, d, x and y sech represents an integer of from 0 to 3, with the proviso that they are not 0 at the same time; e, f, g, h, p, q, r, e, t and us chrepresents an integer of from 0 to 3, with the growth of the same time; e, f, g, h, p, q, r, e, t and us chrepresents an integer of from 0 to 3; and represents an integer of for 0 to 5; and represents an integer of for 0 to 5; and represents an integer of for 0 to 5; and represents an integer of for 0 to 5; and represents an integer of for 0 to 5; and represents an integer of for 0 to 5; and represents and the growth of the same time; e, f, g, h, p, q, r, e, t and us chrepresents and represents and respect to find the same time; e, f, g, h, p, q, r, e, t and us chrepresents and respect to find the same time; e, f, g, h, p, q, r, e, t and us chrepresents and respect to find the same time; e, f, g, h, p, q, r, e, t and us chrepresents and respect to find the same time; e, f, g, h, p, q, r, e, t and us chrepresents and respect to find the same time; e, f, g, h, p, q, r, e, t and us chrepresents and respect to find the same time; e, f, g, h, p, q, r, e, t and us characterised the same time; e, f, g, h, p, q, r, e, t and us characterised the same time; e, f, g, h, p, q, r, e, t and us characterised the same time; e, f, g, h, p, q, r, e, t and us characterised th

Specific examples of the compounds represented by the foregoing general formulae (II) to (IX) include those having the following structural formulae:

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wherein Y<sup>1</sup> to Y<sup>12</sup> each represent a hydrogen atom or a group represented by the general formula (I), with the proviso that at least two of the various compounds have a group represented by the general formula (I). Preferably, Y<sup>1</sup> to Y<sup>12</sup> all are a group represented by the general formula (I).

In the present invention, as the substance which absorbs light to generate heat there may be used any pigment or dye.

As these pigments there may be used commercial pigments and pigments described in Handbook of Color Index (C.I.), "Handbook of Modern Pigments", Japan Pigment Technology Association (1977), "Applied Technology of Modern

ern Pigments", CMC Shuppan (1986), and "Technology of Printing Ink", CMC Shuppan (1984).

Examples of these pigments include black pigment, yellow pigment, orange pigment, brown pigment, red pigment, being pigment, upon pigment, green pigment, thorescent pigment, and polymen-bound dye. Specific examples of the pigments employable herein include isoluble azo pigment, azo lake pigment, cardnersed azo pigment, chielate azo pigment, pithalocyanine pigment, arthraquinone pigment, perione pigment, perione pigment, thioridigo pigment, quinacridone pigment, discazzine pigment, isolndolinone pigment, quinacridone pigment, discazzine pigment, include pigment, discazzine pigment, arthraquinone pigment, arthraquinone, arthr

Such a pigment may be optionally subjected to surface treatment before use. Examples of possible surface treatment methods include a method which comprises coating the surface of the pigment with a resin or wax, a method which comprises attaching a surface active agent to the pigment, and a method which comprises bonding a reactive substance (e.g., siane coupling agent, epoxy compound, polyisocyanate) to the surface of the pigment. These surface treatment methods are turther described in Properties and Application of Metallic Soap', Sachi Shobo (1988). Technology of Infining Ink'; OMC Shuppan (1989), and "Applied Technology of Modern Pigments", OMC Shuppan (1989).

The grain diameter of the pigment is preferably from 0.01 to 10 µm, more preferably from 0.05 µm to 1 µm. As the method for dispersing the pigment there may be used any known dispersion method used in the production of link or loner. Examples of dispersing apparatus which can be employed in these dispersing methods include utilization despersing methods in the despersion of the despersion o

As the dive there may be used any of commercial dyes and known dyes described in references (e.g., "Handbook of Dyes," The Society of synthetic Organic Chemistry, Japan (1970)), Specific examples of dyes armolypable herein include azo dye, metal complex azo dye, prozatione azo dye, anthracquinone dye, phthalocyanine dye, carbonium dye, uninnenimine dye, methins dye, and requirite over

Particularly preferred among these pigments or dyes are those which absorb infrared rays or near infrared rays.

As the pigment which absorbs infrared rays or near infrared rays there may be preferably used caroon black.

As the pigment winch absorbs infrared may or mea immed upsy termed upsy be preceded you be preceded you accord which the barmeter of the dye which absorbs infrared may or near infrared may include cyanine dyes described in JP-A-59-125246, JP-A-59-125266, JP-A-59-1252666, JP-A-59-125266, JP-A-59-125266, JP-A-59-125266, JP-A-59-125266

Further, near infrared absorbents described in U.S. Patent 5,156,938 can be preferably used.

Further, substituted any/benaco(hio)pynyllum salts disclosed in U.S. Patert 3,881,924, trimethinethiapynyllum salts disclosed in JP-A-57-14264 (U.S. Patert 4,327,159), pnyyllum compounds disclosed in JP-A-59-1450153, JP-A-59-220143, JP-A-59-145003, pnyllum salts disclosed in JP-A-59-145001, oyenine dyse disclosed in JP-A-5

Another preferred example of dye is a near infrared absorbing dye represented by the general formula (f) or (II) disclosed in U.S. Patent 4,756,993.

Such a pigment or dye may be incorporated in the image recording material in an amount of form 0.01 to 50% by the preferably form 0.1 to 20% by weight, more preferably form 0.3 to 15% by weight, based on the total solid content in the image recording material. If the content of the pigment or dye falls below 0.01% by weight, no desirable images can be obtained. On the contrary, if the content of the pigment or dye exceeds 50% by weight, the resulting negative-working insour excercion material is and to side on the non-image area during printed.

Such a dye or pigment may be incorporated in the same layer as the phenol derivative or in a layer provided separately of the phenol derivative. If the dye or pigment is incorporated in a separate layer, it is preferably incorporated in a layer adjacent to the layer in which the phenol derivative is incorporated.

The water-insoluble and aquoous alkali-soluble reein to be used in combination with the phenol derivative of the present invention will be further described hereinafter. As the water-insoluble and aquoous alkali-soluble reein there may be used any reein. In particular, a resin having a phenolic hydroxyl group or oldinically unsaturated bond is preferred. Praferred examples of such a reein include the following novolak resins.

Examples of such a novolek resin include cresoftormaldshyde resin, under sea phenollormaldshyde resin, moresoftormaldshyde resin, cresoftormaldshyde resin, moresoftormaldshyde resin, concentration of the more of the season of

resin). In particular, phenolic resins described in JP-A-61-217034 are preferred.

Further, phenoi-modified xylene resins, polyhydroxystyrene, polyhalogenaled hydroxystyrene, and adylfo resins having a phenolic hydroxy group described in JP-A-50-55406, JP-A-51-34711, JP-A-51-36129, JP-A-52-26401, JP-A-62-38454, West German Patents 3,528,390 and 3,528,929, U.S. Patent 4,724,195, JP-A-5-230139, JP-A-5-230140, and JP-A-7-333389 may be used.

Examples of resin having an olefinically unsaturated bood which can be preferably used in the present invention include those described in JP-B-3-63740, U.S. Pateria \$,376,138 and 3,556,793, JP-A-52-988, and JP-B-60-37123. The resin having an olefinically unsaturated bond preferably contains an alkali-soluble monomer as a copplymerizable component to enhance the developability of the negative-working image recording material in an aqueous alkaline classification.

As the alkali-soluble group there may be used an acidic group having a pKa value of not more than 14. Examples of such an acidic group include -SO<sub>3</sub>H, -OP(O)(OH)<sub>2</sub>, -P(O)(OH)<sub>2</sub>, -COOH, -COOH, -COOHSO<sub>2</sub>, -COOHSO<sub>2</sub>, -SO<sub>2</sub>NH-, and phenolic OH group.

As the aqueous alkali-soluble resin there is preferably used one having a weight-average molecular weight of from 500 to 400,000 and a number-average molecular weight of from 200 to 150,000.

These aqueous alkali-soluble resins may be used singly or in combination. The amount of such an aqueous alkalisoluble resin to be incorporated is from 5 to 99% by weight, prefeably from 30 to 99% by weight based on the total weight of the photosensitive composition as calculated in terms of solid content.

The image recording material of the present invention may optionally further comprise various additives incorpo-

For example, a multifunctional monomer having two or more radically polymerizable ethylenic double bonds per molecule may be incorporated in the image recording material layer. Examples of such a compound include striptene glycol differnitacytiste, eighthere glycol differnitacytiste, brane did differnitacytate, trimethylorylate, trimethylorylate, trimethylorylate, trimethylorylate, trimethylorylate, trimethylorylate, trimethylorylate, trimethylorylate, and trifernitacytate dispersatesythriated and trimethylorylate dispersatesythriated and trimethylorylated and trimethylorylated

Further examples of actifives which can be incorporated in the image recording material layer include an ally/lether for improving coatability (e.g., ethyl cellubose, metry cellubose), a surface active agent (e.g., fluorine surface active agent), and plasticate for providing film softening property and abrasion resistance (e.g., tricresy) phosphate, directly phosphate, tributyl phosphate, tributyl citrate, polyethylene glycol, polypropylene glycol. The amount of these additions to be incorporated depends on their purpose but is normally from 0.5 to 30% by weight beased on the total sold content in the image recording material.

A representative example of printing-out agent for providing a visible image immediately after the heat generation due to exposure is a combination of a compound which releases an acid when heated due to excouse and an organic object agent of forming a sait. Specific examples of such a printing-out agent employable herein include a combination of heated and printing-out agent employable herein include a combination of heated and as assit-forming organic dye as disclosed in JP-A-53-38209 and JP

The printing-out agent and dive are incorporated in the image recording material in an amount of from 0 to 30% by
46 weight.

If the substance of the present invention which absorbs light to generate heat is used to obtain a visible image having a sufficient density, such a dye doesn't need to be added. For the purpose of accelerating crosslinking reaction, only a printing-out agent (adid operator) may be added.

The image recording material of the present invention may be provided by coating a support with the toregoing components in the form of solution or dispersion in a solvent. Examples of the solvent employable herein include mather and, eithand, isopropyl alcohol, a-butyl alcohol, a-butyl alcohol, ethylene dichloride, cyclohaxanone, acetone, methyl ethyl ketone, ethylene glycol, ethylene glycol monosethyl ether, Z-metrovycht sociata, 1-methyny2-propend, 1-methoxy2-properd, acetate, N-metherythormamice, N-Morinertylecatamide, tettalydrofurar, dioxane, dimethyl sulfoxide, ethyl acetate, methyl lactate, and ethyl lactate. These solvents may be used eingly or in admixture.

A mixed solvent obtained by adding to such a solvent or mixed solvent a small amount of water or a solvent which doesn't dissolve phenol derivatives or high-molecular compounds therein, such as toluene, is preferred, too.

The concentration of the foregoing components (solid content) in the solvent is from 1 to 50% by weight.

If the solution or dispersion thus obtained is coated and dried, drying is preferably effected at a temperature of from

50°C to 120°C. The drying process may comprise predrying at a low temperature and subsequent drying at a higher temperature. Alternatively, drying may be effected at high temperature if the solvent and concentration are properly selected, though drying at 150°C or higher is not desirable due to the heat-sensitive recording material.

The applied amount of the costing meterical depends on the purpose. If the negative-working image recording meterial is used o<sub>8</sub>, as a protosensitive lithography in printing plate, the enerother lithography printing plate, the enerother lithography printing plate), the applied amount of the costing meterial is, the higher is the sensitivity but the porcer are physical properties of photosensitive layer. If necessary, a matter or matte layer may be provided on the photosensitive layer. Further, an undercosting layer may also be provided on the photosensitive layer.

In the present place of the support to be coated with an image recording material of the present invention include paper, appare trainisect with a plastic (e.g., polyetyfriene, polytagreine, place made of a meet acut as aluminum (including aluminum silety), zinc and capper, tim made of a plastic such as callulose capporate, cellulose accentance, cellulose accentance, cellulose proprients, cellulose proprients, cellulose accentance, cellulose, cell

The support having a metallic surface, particularly aluminum surface, is preferably subjected to a proper hydrophilic

The hydrophilic treatment may be conducted as follows. For example, the surface of an aluminum pixthe is grained by a mechanical graining method such as wire brush graining, with brush graining with a surry of barester grains and ball graining, a chemical graining with HF. ACIs or HCI as an etchant, an electolytic graining with third acid or hydrochloric acids as an electrolyte or a composite thereoty optionally etched with an acid or a fisalit, and then anodically ordered with DC or ACI current in suffuris acid, phosphoric acid, oxatic acid, boric acid, chromic acid, suifamic acid or motive thereof to form a rigid assessed limit thereon.

Such a passive film can render the aluminum surface bydrophile. If necessary, the aluminum surface thus treated is preferably subjected to breatment with silicate (e.g., sodium silicate, potassium silicate) as disclosed in U.S. Patent 3.214,065 and 3.181.461, freatment with potassium fluorocirconate as disclosed in U.S. Patent 3.291.247, treatment with abyl titrante as disclosed in British Patent 1,186.599, treatment with polyacrylic acid as disclosed in German Patent 1,091.439, treatment with polyacrylic acid as disclosed in German Patent 1,200.447, treatment with phylosylic acid as disclosed in U.S. Patent 3.309.591, composite acid as disclosed in JP-8-44-699. Terestment with phylosylic acid as disclosed in U.S. Patent 3.309.591, composite treatment with as a hydrophilic organic high-molecular compound and a divalent metal as disclosed in JP-A-68-16993 and JP-A-68-1891, or undercoating with a water-soluble polymer having a sulforic group as disclosed in JP-A-69-16993 roll of the disclosed in JP-A-69-16993 roll of t

As the active my source for use in imagewise exposure there may be used a mercury vapor lamp, metal hailds image, zenoi lamp, chemical lamp, carbon-are lamp or be life. Examples of radiation include electron reys, X rays, kn beam, and far infrared rays. Further, g-line, Line, Deep-UV rays, and high dentity energy beam (laser beam) may be used. Examples of such a laser beam include helium-neon laser, rayon laser, laybun see, helium-cadmini laser, and K/F exter slaser. In the present invention, a light source which entits light in the range of near infrared to intrand is desirable. In outcidar, a solid leaver or semiconductor laser is preferred.

46 As the developer for the negative-working image recording material according to the present invention or its replenisher there may be used any forom acqueus allesine soution. Examples of such an alkali includie inorganic alkali sales auch as ecdium elicate, podasium silicate, sodium teriary phosphate, potassium teriary phosphate, ecdium carbonate, potassium secondary phosphate, potassium teriary phosphate, ecdium toriary phosphate, endit in experimental potassium experiments according phosphate, ammonium hydrogencarbonate, sodium toriate, sodium toriate, ammonium borate, sodium toriate, sodium toriate, ammonium toriate, ammonium toriate, sodium toriate, ammonium toriate, sodium toriate, sodium toriate, ammonium toriate, sodium toriate, sodium toriate, ammonium toriate, ammonium toriate, sodium toriate

The developer or its replenisher may optionally comprise various surface active agents or organic solvents incorporated therein for the purpose of accelerating or inhibiting developability and enhancing dispersibility of development residue and ink-receptivity of the image area on the printing plate. Preferred examples of such surface active agents include anionic, cationic, nonionic and amphotetic surface active agents.

Further, the developer or its replenisher may optionally comprise a reducing agent such as hydroquinone, resorcin and sodium and potassium sait of inorganic acid such as suffutous acid and hydrogensulfurous acid, organic carboxylic acid, defoaming agent, hard water soffener, etc. incorporated therein.

Preferred examples of the developer employable herein Include those described in JP-A-54-62004 and JP-B-57-7427, a developer composition comprising observal elochol, an anionic surface active agent, an alvains agent and water as disclosed in JP-A-51-77401, a developer composition comprising an appeass solution containing benzyl alcohol, an anionic surface active agent and a water-soluble sulfurous acid as disclosed in JP-A-53-44202, and a developer composition comprising an organic schemt having a solubility to water of not more than 10% by weight at normal temperature, an alkaline agent and water as disclosed in JP-A-53-155555.

in the case where the image recording material according to the present invention is used as a printing clate, the invention is present in the present of the present invention is used as a printing clate the repletisher is prelensibly subjected to post-treatment with a trinsing solution containing a washing water and a surface active agent or a desensitizing solution containing gum arabic or a stand derivative. The post-treatment may comprise these treatments in combristion.

In the recent plate-making and printing industry, automatic developing machines for printing plate have been widely used from the standpoint of the reticentization and standardization of plate making process. These automatic developing machines normally consist of a development zone and a post-treatment zone. These automatic developing machines comprise an apparatus for carrying a printing plate, various processing tanks, and a spraying apparatus. In operation, various processing solutions which have been pumped up are oprayed through the respective nozzle onto see an exposed printing plate withort is being carried byte dwith a processing solution which being carried guided by a ucharreged guide rol. In such an automatic processing process, the various processing solutions may be replenished by its replenisher deporting on the throughout covariant lime, etc.

Further, a so-called throwaway system which comprises processing with a substantially unused processing solution may be employed.

The present invention will be further described in the following examples, but the present invention should not be construed as being limited thereto.

# SYNTHESIS EXAMPLE 1-1 (Synthesis a of Compound (X))

2 g of 1-(a-methyl-a-(4-hydroxyphenyf)ethyl)-4-(a,a-bis(4-hydroxyphenyf)ethyl)benzene (compound represented by the general formula (X) wherein Y is Y are a hydrogen atom at the same time) was dissolved in 100 mt of a 10% aquecus solution of potassium hydroxida. To this reaction solution was terne added cloowise 60 mt of 27% formalin with stirring at room temperature in 1 hour. The reaction solution was further stirred at room temperature for 6 hours, and 35 then poured into an aqueous solution of sufficient of undergo crystallization. The paste precipitate thus obtained was thoroughly washed with welat-, and then recrystallized from 30 mt of methanol to obtain a white powder (yeld: 20 g).

The compound thus obtained was then identified as a hexamethylolated product of 1-[-methyl-c-(-hydroxypte-ny) ethyl-4-(-hydroxypte-ny) ethyl-4-(-h

# SYNTHESIS EXAMPLE 1-2 (Synthesis b of Compound (XI)) 20 g of the hexamethylolated product obtained in

Synthesis Example 1-1 (compound represented by the general formula (X) wherein Y¹ to Y<sup>6</sup> are a methylol group at the same time) was dissolved in 1,000 m² of methanol under heating. To the solution was then added 1 m² of concentrated suffuria acid. The reaction solution was then heated under reflux for 12 hours. The reaction solution was then allowed to coci. To the reaction solution was then added 2 g of potassium carbonate. The reaction mixture was stirred, and then concentrated. To the reaction solution was then added 30 m² of athyl accetate. The reaction solution was washed with water, and then dried. The solvent was then distilled off to obtain a white solid (yied: 22 g).

The compound thus obtained was then identified as a hexamethosymethylated product of 1-(a-methyl-cu-(4-hydroxyyn-my)derhyl-4-(a--bis(4-hydroxy-hyny)derhylbenzene (compound represented by the general formula (X) wherein Y to Y's are a methosymethyl group at the same time) by NMR. The purity of the hexamethosymethylated product was 90% as determined by reversed phase HPLC (column: Shimpac CLC-ODS (Shimadzu Corp.); solvent: methanol/water si (mixing ratio = 60M40 50Y10 by volume)).

### SYNTHESIS EXAMPLE 1-3 (Synthesis c of Compound (X))

20 g of the hexamethylolated product obtained in Synthesis Example 1-1 (compound represented by the general

tormula (X) wherein Y is Y are a methylol group at the same time) was dissolved in 1,000 mf of sharol under heating. To the solution was then added if not of concentrated suffucia and The reaction solution was then added in ordinary to 12 hours. The reaction solution was then allowed to cool. To the reaction solution was then added 2 g of potassium corthoals. The reaction motivare was stirred, and then concentrated. To the reaction solution was then added 300 mf of ethyl acetate. The reaction solution was washed with water, and then dried. The solvent was then distilled off to obtain a colories soll win thater (videt 2.2 g.).

The purity of the hexaethoxymethylated product (compound represented by the general formula (X) wherein Y<sup>1</sup> to Y<sup>6</sup> are an ethoxymethyl group at the same time) was 70% as determined by reversed phase HPLC (column: Shimpac CLC-ODS (Shimadu Corp.) solvent: methanolwater (mixing ratio = 60/40 to 90/10 by volume).

#### SYNTHESIS EXAMPLE 1-4 (Synthesis d of Compound (X))

20 g of the hexamethylolated product obtained in Synthesis Example 1-1 (compound represented by the general formula (f) wherein Y <sup>1</sup> to Y <sup>2</sup> are a methylol group at the same time) was dissolved in 200 mt/ of 1-methoxy-2-propool under heating. To the solution was then actived 1 mt/ of concentrated suffuric acid. The reaction solution was then heating under reflux for 12 hours. The reaction solution was then allowed to cool. To the reaction solution was then added 2 g of potasytum carbonate. The reaction mixture was stirred, and then concentrated. To the reaction solution was then added 300 mt/ of ethyl acetate. The reaction solution was washed with water, and then dried. The solvent was then distilled of the obtain a podinces of low matter (visit 13 a).

The content of the hexaetherfiled product (compound represented by the general formula (X) wherein Y<sup>1</sup> to Y<sup>5</sup> are a 2-methoxy-t-methylethoxymethyl group at the same time) was 55% as determined by reversed phase HPLC (column: Shirmack CLC-ODS (Shirmack) CCADS (Shirmack) CCADS (Shirmack) CCADS (Shirmack) (CADS (Shirmack) (CADS (Shirmack)) (CADS (Shirma

# SYNTHESIS EXAMPLE 1-5 (Synthesis e of Compound (X))

20 g of the hexamethylolated product obtained in Synthesis Example 1-1 (compound represented by the general formula (X) wherein V to V<sup>6</sup> are a methylol group at the same time) was dissolved in 100 mf of methylene chloride under heating. To the solution was then added 10 mf of acetyl chloride. The reaction mixture was then stirred at a terms of the contractive of 25°C for 12 hours. The reaction solution was weshed with water, and then dried. The solvent was then dissilled off to obtain a coloriess of im matter (vield 21 a).

The compound thus obtained was then identified as a hexakisacetoxynethylated product of 1-(a-methyl-cq-4-hydroxyphenylpel-fq,a-chis(4-hydroxyphenylpel-fq)ebplenzene (compound represented by the general formula (X) wherein's '0' are an accionymethyl group at the same time) by MRN. The purity of the hexakisacetoxymethylated wherein's '0' are an accionymethylated by MRN. The purity of the hexakisacetoxymethylated by State of the compound of the same time) by MRN. The purity of the hexakisacetoxymethylated by MRN.

### SYNTHESIS EXAMPLE 2-1 (Synthesis a of Compound (XI))

17.3 g of p.c.i.c.\*-Insig4-hydroxyphenyl\(0.1.3.5\)-frileopropylbenzene (compound represented by the general formula (XI) wherein' 1 to 7 are a hydrogen atom at the same time) was dissolved in 80 g of a 10% acqueous solution of sodium hydroxide. To the reaction solution was then added 15 g of methanol. To the reaction solution was then added of powers 30 g of 37% formalin. After the completion of the drophes addition, the reaction solution was theat added dropwise 30 g of 37% formalin. After the completion of the drophes addition. The reaction solution was theat cystalized form an expectation of a certification of the drophes addition. The reaction solution was the crystalized form an expectation of a certification of the drophes addition. The reaction solution was the crystalized form and the drophes it was the manufacture of the complete addition of the certification of the drophes addition of the complete addition of the certification of the drophes addition. The reaction solution was heated to a temperature of 4 of "O" where a function of the certification of the drophes addition. The reaction solution was heated to a temperature of the certification of the certi

The compound thus obtained was then identified as a hexamethylolated product of  $\alpha$ ,  $\alpha$ ( $\alpha$ '-tris(4-hydroxyphenyl)-1,3,5-triisopropylbenzene (compound represented by the general formula (XI) wherein Y I by Y are a methylol group at the same time) by NMR. The purity of the hexamethylolated product was 90% as determined by reversed phase HPLC (column: Shimpac CLC-ODS (Shimadzu Corp.); solvent: methanol/water (mixing ratio = 60/40 to 90/10 by volume)).

### SYNTHESIS EXAMPLE 2-2 (Synthesis b of Compound (XI))

20 g of the hexamethylolated product chalance in Synthesis Example 2-1 (compound represented by the general formula (XI) wherein Y<sup>1</sup> to Y<sup>6</sup> are a methylol group at the same time) was dissolved in 1,000 m of or intellunal under heatest ing. To the solution was then added 1 m of concentrated sulturic acid. The reaction solution was then heated under reflux for 9 hours. The reaction solution was then allowed to cool. To the reaction solution was then added 2 g of potassium carbonate. The reaction mixture was stirred, and then concentrated. To the reaction solution was then added 30 m of of ethyl acetate. The reaction solution was washed with water, and then diried. The solvent was then distilled off to obtain a colorless of winterfor videl: 22 c).

The purity of the hexakismethoxymethylated product (compound represented by the general formula (XI) wherein Ye are a methoxymethyl group at the same time) was 85% as determined by reversed phase HPLC (column: Shinnac CLC-ODS (Shimadz) Corp.); solvent methanol/water (mixing ratio = 60/4 reversed phase HPLC (column: Shinnac CLC-ODS (Shimadz) (Corp.); solvent methanol/water (mixing ratio = 60/4 reversed phase HPLC (column: )

#### SYNTHESIS EXAMPLE 3-1 (Synthesis a of Compound (XII))

10 g of ca.c.'c'-tetralside/-hydroxyphenyl-p-vylene (compound represented by the general formula (XII) wherein '10 ye' are a hydrogen attom at the same time) was dissolved in 25 ml of a 15% aqueous solution of solution To the reaction solution vas then added 15 ml of methanol. To the reaction solution was then added dropwise 20 g of 37% formalin. After the completion of the dropwise addition, the reaction solution was therefore to empletion of the dropwise addition, the reaction solution was therefore the empletion of effect of the solution and the solution was the recompletion of the dropwise 20 g of 40 where it was then allowed to undergo reaction for 12 hours. The reaction solution was then crystallized from an aqueous solution of acetic acid. The solid thus obtained was thoroughly washed with water, and then resturried with 100 m/c of ethyl acetate to obtain a white power (yelds: 11.4).

### 20 SYNTHESIS EXAMPLE 3-2 (Synthesis b of Compound (XII))

20 g of the octamentylolated product obtained in Synthesis Example 3-1 (compound represented by the general formula (XII) wherein Y 1 to Y<sup>2</sup> are a methylol group at the same time) was dissolved in 1,000 m² of methanol under heading. To the solution was then added 1 m² of concentrated sulfuric acid. The reaction solution was then heated under are reflux for 10 hours. The reaction solution was then allowed to cool. To the reaction solution was then added 2 g of potassium carbonate. The reaction mixture was stirred, and then concentrated. To the reaction solution was then added 300 m² of ethyl acetate. The reaction solution was washed with water, and then diried. The solvent was then dissilled off to obtain a light yellow oily imstertly (righed: 21 g).

The purity of the octal/smethoxymethylated product (compound represented by the general formula (XVI) wherein Y<sup>1</sup> to Y<sup>2</sup> are a methoxymethyl group at the same time) was 67% as determined by reversed phase HPLC (column: Shimpac (C-ODS (Shimadzu Corx); solvent: methanol/water (mising ratio = 60/40 to 90/10 by volume)).

### SYNTHESIS EXAMPLE 3-3 (Synthesis c of Compound (XII))

39 20 g of the octamethyloteted product obtained in Synthesis Example 3-1 (compound represented by the general formula (XII) wherein Y 1o Y<sup>2</sup> are a methylot group at the same time) was dissolved in 800 m² of ethanol under heating. To the solution was then added 1 m² of concentrated sulfuric add. The reaction solution was then heated under reflux for 12 hours. The reaction solution was then allowed to cool. To the reaction solution was then heated under reflux carbonats. The reaction instruct was stirred, and then concentrated. To the reaction solution was then added 20 m of earthy acetates. The reaction solution was then added 300 m of earthy acetates. The reaction solution was then added 300 m of earthy acetates. The reaction solution was washed with water, and then dried. The solvent was then distilled off to obtain a copinges of matter (viet 23 o.).

The purity of the octakisethoxymethylated product (compound represented by the general formula (XVI) wherein Y<sup>1</sup> to Y are an ethoxymethyl group at the same time) was 72% as determined by reserved phase HPLC (column: Shimnac CLC-ODS (Shimnac) cont.), solvert: methano/water (mixing ratio = 60/40 to 90/10 by volume).

### SYNTHESIS EXAMPLE 3-4 (Synthesis d of Compound (XII))

20 g of the octamethylotated product obtained in Synthesis Example 3-1 (compound represented by the general formula (XII) wherein Y1 to Y<sup>3</sup> are a methylot group at the same time) was dissolved in 100 m/ of methylene chloride our drief health of the solution was then added to Im of diseably offloride. The reaction solution was then stirred at a temperature of 25°C for 12 hours. The reaction solution was washed with water, and then dried. The solvent was then distilled off to obtain a colories of im matter (rief 22 d).

The purity of the octal/sacetoxymethylated product (compound represented by the general formula (XII) wherein Y<sup>1</sup> to Y<sup>2</sup> are an acetoxymethyl (proug at the same time) was 75% as determined by reversed phase HPLC (column: 55 Shimpac CLC-ODS (Shimadzu Corp.): solvent: methanolylater (mixing ratio = 60/40 to 90/10 by volume)).

# SYNTHESIS EXAMPLE 4-1 (Synthesis a of Compound (XIII))

11 g of 1,3,3,5-tetrakis(4-hydroxyphenyl)-pentane (compound represented by the general formula (XIII) wherein Y1

to V<sup>2</sup> are a hydrogen atom at the same time) was dissolved in 40 g of a 10% aqueous solution of sodium hydroxide. To this reaction solution was then actived 10 g of methanol. To the reaction mixture was then actied dropwise 2g of 37% formain. After the completion of the dropwise addition, the reaction solution was heated to a temperature of 40°C where it was then further allower to undergo reaction for 12 hours. The reaction solution was then crystalized from an equeous solution of acetic acid to obtain a white viscous solid. The solid thus obtained was thoroughly washed with water, and then dried at room temperature under reduced pressure (yield: 13.6 g).

The compound thus obtained was then identified as an octamethylotated product of 1,3,3,5-tetrakis(4-hydroxyphe-nyl)-pantane (compound represented by the general formula (XIII) wherein Y 1 to Y<sup>2</sup> are a methylot group at the same time) by NMR. The purity of the octamethylotated product was 93% as determined by reversed phase HPLG (column: 95 Shirmpac CLG-ODS (Shirmadau Corp.); solvent: methanol/water (mixing ratio = 60/40 to 90/10 by volume).

# SYNTHESIS EXAMPLE 4-2 (Synthesis b of Compound (XIII))

20 g of the octamethylolated product obtained in Synthesis Example 4-1 (compound represented by the general formula (XIII) wherein Y' to Y' are a methylol group at the same time) was dissolved in 1,000 mt of methanol under heading. To the solution was then acided 1 mt of concentrated suthizin add. The reaction solution was then heated under reflux for 10 hours. The reaction solution was then allowed to cool. To the reaction solution was then added 2 g of potassium automates. The reaction institute was stitred, and then concentrated. To the reaction solution was then added 30 of of eithyl acetate. The reaction solution was washed with water, and then orderated on the solvent was then distilled off to obtain a joint vallew of with water (ridel: 21 g).

The purity of the octakismethoxymethylated product (compound represented by the general formula (XIII) wherein Yo are a methoxymethyl group at the same timely was 85% as determined by reversed phase HPLC (column: Shimpac OLC-ODS (Shimaczu Corp.); solvent methanol/water (mixing ratio - 60/40 to 90/10 by volume)).

### 85 SYNTHESIS EXAMPLE 5-1 (Synthesis a of Compound (XVI))

The compound thus obtained was then identified as a dodecamethylolated product of  $\alpha, \alpha, \alpha', \alpha', \alpha''$  an identified as a dodecamethylolated product of  $\alpha, \alpha, \alpha', \alpha''$  an invariant hydroxyphany)-1,3,5-triethylobanzene (compound represented by the general formula (XVI) wherein Y  $^{1}$  by  $^{1}$  are a methylol group at the same time) by NMR. The purity of the dodecamethylolated product was 80% as determined by reversed phase HPLC (column: Shimpac CLC-ODS (Shimadzu Corp.); solvent: methenoliwater (nixing ratio = 80/40 to 90/10 by volume).

# 40 SYNTHESIS EXAMPLE 5-2 (Synthesis b of Compound (XVI))

23 g of the dode-amethylosited product obtained in Synthesis Example 5-1 (compound increasanted by the general formula (V/I) wherein Y<sup>1</sup> to Y<sup>12</sup> are a methylol group at the same time) was dissolved in 1,000 m of or methanol under heating. To the solution was then aided 1 m of concentrated safulure and. The reaction solution was then heated under a reflux for 13 hours. The reaction solution was then aideved to cool. To the reaction solution was then added 20 g of polar-situm carbonate. The reaction mixture was stirred, and then concentrated. To the reaction solution was then added 30 m of of ethyl acetate. The reaction solution was washed with water, and then dired. The solvent was then distilled off to obtain a joint yellow only matter (yield: 21 g).

The purity of the dodecakismethoxymethylated product (compound represented by the general formula (XVI) when in Y to Y<sup>12</sup> as methoxymethyl group at the same time) was 75% as determined by reversed phase HPLC (column: Shimpac CLC-ODS (Shimsatu Corp.); solvent: methanol/water (mixing ratio = 60/40 to 90/10 by volume).

#### [Preparation of substrate]

A 0.3-mm thick alumnum plate (quality: 1050) was solvent-cleaned so that it was depreased, prained with a rylon or and an aqueous suspension of 400-mesh pumice, and then thoroughly washed with vater. The alumnum plate thus grained was dipped in a 25% aqueous solution of sodium hydrodde at a temperature of 45°C for 9 seconds so that it was exthed, washed with water, dipped in a 20% initio acid for 20 seconds, and then washed with water. The ethical amount of the grained surface of the alumnum plate was shout 3 gift. The alumnum plate thes treated was subjected.

to oxidation with a DC current in a 7% sulfuric acid as an electrotyte at a current density of 15 Actm<sup>2</sup> to obtain an oxidized film having a density of 3 g/m<sup>2</sup>, washed with water, and then dried. The aluminum plate thus oxidized was costed with the following undercoating solution, and then dried at a temperature of 80°C for 30 seconds. The density of the undercoating layer after dried was 10 mg/m<sup>2</sup>.

(Undercoating solution)	
β-Alanine	0.1 g
Phenylphosphonic acid	0.05 g
Methanol	40 g
Pure water	60 g

[Preparation of carbon black dispersion)

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The following components in the following weight proportion were subjected to dispersion with glass beads for 10 minutes to obtain a carbon black dispersion.

	Carbon black	1 part by weight
	Copolymer of benzyl methacrylate and methacrylic acid (molar ratio: 71 : 29; weight-average molecular weight: 70,000)	1.6 parts by weight
	Cyclohexane	1.6 parts by weight
,	Methoxypropyl acetate	3.8 parts by weight

[Preparation of negative-working lithographic printing plate]

### EXAMPLES 1 TO 13 AND COMPARATIVE EXAMPLES 1 TO 4

The aluminum plate thus obtained was coated with the following photosensitive layer, and then dried at a temperature of 100°C for 2 minutes to obtain a negative-working photosensitive lithographic printing plate. The density after dried was 2.0 d/m<sup>2</sup>.

	Photosensitive layer	
45	Carbon black dispersion mentioned above	2.4 g
**	Phenol derivative	(as set forth in Table 1)
	Phenol-formaldehyde novolak (weight-average molecular weight: 12,000)	(as set forth in Table 1)
	4-(p-N,N-bis(ethoxycarbonylmethyl) aminophenyl-2,6-bis(trichloromethyl) -S-triazine	0.02 g*1
50	Megafac F-176 (fluorinic surface active agent available from Dainippon Ink & Chemicals, Inc.)	0.06 g
	Methyl ethyl ketone	15 g
	2-Methoxy-1-propanol	12 g

(\*1: Not added in Examples 1 and 2)

.

	Used amount		۵.	neno!	Phenol derivative	ive	1	-	Printable
	of	General						Used	number
	novolak (g)	formula	S	vnthe	Synthesis example	ple	1	amount	of sheets
Example 1	2.10	(x)	Compound of Synthesis Example	of Sy	nthesis	Example	1-1	0.21	30,000
Fyample 2	2.00	×	Compound of Synthesis Example	of Sy	nthesis	Example	1-2	0.21	25,000
Example 3	2.10	(x)	Compound	of Sy	nthesis	Synthesis Example	1-3	0.20	30,000
Example 4	2.05	3	Compound of	of	nthesis	Synthesis Example	1-4	0.21	20,000
Example 5	2.00	æ	Compound	of Sy	nthesis	Synthesis Example	1-5	0.22	35,000
Example 6	2.07	(XI)		ф	nthesis	Synthesis Example	2-1	0.21	30,000
Example 7	2.07	(XI)	Compound	οţ	nthesis	Synthesis Example	2-3	0.21	30,000
Example 8	2.11	(XIX)	Compound	οĘ	nthesis	Synthesis Example	3-1	0.21	30,000
Example 9	2.11	(XII)	Compound of		nthesis	Synthesis Example	3-2	0.23	25,000
Example 10	2.10	(XII)	Compound of	of Sy	nthesis	Synthesis Example	3-3	0.21	30,000
Example 11	2.08	(XIII)	Compound of	of Sy	nthesis	Synthesis Example	4-1	0.21	25,000
Example 12	2.08	(XIII)	Compound of	of Sy	nthesis	Synthesis Example	4-2	0.21	25,000
Example 13	2.05	(XAI)	Compound of Synthesis Example 5-1	of Sy	nthesis	Example	5-1	0.20	30,000
Comparative								Not	Image not
Example 1	2.10	1		1				added	formed
Comparative			Starting material of	mater	ial of				Image not
Example 2	2.09	(x)	Synthesis Example	Bxam	ple 1			0.20	formed
Comparative									Image not
Example 3	2.12	(XXI)		1				0.22	formed
Comparative									
Example 4	2.10	(XXII)		1	-		-	93.6	10,000

The compound used in Comparative Example 2 was a compound represented by the general formula (X) wherein Y to V<sup>3</sup> are a hydrogen atomat the same time. The compounds (XXI) and (XXII) used in Comparative Examples 3 and 4 have the following structures having one and two benzene nuclei, respectively. These compounds are described in JP-B-1-49832.

$$HOCH_2$$
 $CH_3$ 
 $HOCH_2$ 
 $CH_3$ 
 $CH_2OH$ 
 $CH_2OH$ 
 $CH_2OH$ 

The negative-vorking photosensitive thiographic printing plate thus obtained was exposed to light beam from a VAG base which had been adjusted such that the output thereof was 700 mW on the surface of the printing plate, and then processed through an automatic processor filled with a developer DP-4 (1 : 8) available from Full Photo Film Co., Ltd. and a rivering solution FP-8 (1 : 7) available from Full Photo Film Co., Ltd. end a rivering solution FP-8 (1 : 7) available from Full Photo Film Co., Ltd. obtain a negative inegar. The lithough printing plate was then used in printing by a Hidel SOR+K2 printer. The results are set forth in Table 1. Examples to 10 :3, in which the phenol derivatives of the present invention were used, gave good printed matters. On the contray, 45 the comparative examples, free of phenol derivatives of the present invention, could form no image. In Comparative Example 4, which formed an image, the printing-the number of sheets was small.

### COMPARATIVE EXAMPLE 5

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The procedure of Example 1 was followed to prepare a negative-working photosensitive lithographic printing plate except that the photosensitive solution was prepared free of carbon black dispersion. The photosensitive lithographic printing plate thus prepared was exposed to light, and then developed in the same manner as in Example 1. As a result, the photosensitive film was totally dissolved in the developer. Thus, no image was obtained.

## 55 EXAMPLES 14 TO 19 AND COMPARATIVE EXAMPLES 6 AND 7

The same aluminum plate as used in Examples 1 to 13 was coated with the following photosensitive layer, and then did at temperature of 100°C for 2 minutes to obtain a negative-working photosensitive lithographic printing plate. The density after direl days 2.0 g/m².

# Photosensitive layer

Carbon black dispersion mentioned above	2.2 g
Phenol compound of Synthesis Example 1-2	0.25 g
Alkali-soluble resin mentioned below	2.05 g
Megafac F-176 (fluorinic surface active agent available from Dainippon Ink & Chemicals, Inc.)	0.06 g
Methyl ethyl ketone	15 g
2-Methoxy-1-propanol	12 g

Alkali-soluble resin

# EXAMPLE 14

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16

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(-cH2-CH-)-OOC-}-OH

EXAMPLE 15

EXAMPLE 16

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EXAMPLE 17

$$\begin{array}{c} \left( \text{CH}_{2} \overset{\text{CH}_{3}}{\leftarrow} \right)^{25} \\ \text{CONHSO}_{2} & \left( \text{CH}_{3} \overset{\text{CH}_{2}}{\leftarrow} \right)^{10} \\ \text{COO} & \left( \text{COO} \overset{\text{COO}}{\leftarrow} \right)^{10} \\ \text{COO} & \left( \text{CH}_{2} \overset{\text{CH}_{3}}{\leftarrow} \right)^{10} \\ \text{COO} & \left( \text{CH}_{2} \overset{\text{CH}_$$

EXAMPLE 18

EXAMPLE 19

$$\left( \begin{array}{c} CH_2 \stackrel{C}{C} \stackrel{C}{C} \stackrel{G}{C} \\ \stackrel{C}{C} \stackrel{G}{O} \stackrel{G}{O} \end{array} \right)^{83} \left( \begin{array}{c} CH_2 \stackrel{C}{C} \stackrel{G}{C} \stackrel{G}{C} \\ \stackrel{C}{C} \stackrel{G}{O} \stackrel{G}{O} \\ \end{array} \right)^{17} \quad .$$

#### COMPARATIVE EXAMPLE 6

#### COMPARATIVE EXAMPLE 7

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(Composition ratio is represented by mol)

The negative-working photosensitive litrographic printing plate thus obtained was exposed to light beam from a VAG laser which had been adjusted such that the output thereof was TOO mill wo me be unable of the printing plate, and then developed with an expense solution containing sodium carbonate and sodium hydrogencationate. As a result, Examples 14 to 19, In which an acytofic resin having phenol ring or all tygroup was used, give an engative image. On the contary, in Comparative Examples 6 and 7, in which an acrylic resin free of phenol ring or ally group was used, since the photogensitive film was totally dissolved in the developer, and no image was thus obtained.

### EXAMPLES 20 TO 22 AND COMPARATIVE EXAMPLES 8 AND 9

A photosensitive layer which had comprised a dye set forth in Table 2 instead of the carbon black dispersion used in Example 1 was applied to the substrate, and then dried in the same manner as in Example 1 to obtain a negativeworking photosensitive lithographic printing plate.

Photosensitive solution	
Dye set forth in Table 2	0.2 g
Phenol compound of Synthesis Example 1-2	(as set forth in Table 2)
Phenol-formaldehyde novolak (weight-average molecular weight: 12,000)	(as set forth in Table 2)
4-(p-N,N-bis(ethoxycarbonylmethyl) aminophenyl)-2,6-bis(trichloromethyl) -S-triazine	0.02 g*2
Megafac F-176 (fluorinic surface active agent available from Dainippon Ink & Chemicals, Inc.)	0.06 g
Methyl ethyl ketone	15 g
2-Methoxy-1-propanol	12 g

(2: Not added in Example 21)

Table 2

	Dye	Used amount of pheno compound (q)	l Printable number of sheets
EXAMPLE 20	1	0.35	25,000
EXAMPLE 21	1	0.34	20,000
EXAMPLE 22	2	0.35	25,000
Comparative Example 8	Not ad	ded 0.34	Image not formed
Comparative Example 9	1	Not added	Image not formed

Dve 1

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Dye 2

$$\begin{array}{c|c} S \\ CH^2 CH^2 CH^2 CH CH = \begin{array}{c} S \\ CH CH CH \\ C_2 H_5 \end{array}$$

The negative working photocenetive lithographic printing plate thus obtained was exposed to light from a semiconductor laser (wavelength: 825 mrs. spot dismatter 14e<sup>2</sup> = 11.9 mr) which had been explicated such that the objupt thereof see was 110 mW on the autrace of the printing plate at a linear speed of 8 mVsc. The printing plate hus exposed was then developed in the same manner as in Example 1. As a result, in Examples 20 to 22, a fine line hexing a width of 10 mW was formed. The lithographic printing plate thus obtained was then used to print on a high quality paper with a commercial ink by a Type SOR-K2 printer available from Hideberger Druckmaschinen Aktiengeselechaft. The results are set forth in Table 2. Examples 20 to 22, in which the planol delarative of the present invention was used in combination with set along called the planol delarative Example 8, which was free of dye, and Comparative Example 9, which was free of dye, and Comparative Example 9, which was free of dye, and Comparative Example 9, which was free of dye, and Comparative Example 9, which was free of dyen of the present invention, formed no Image.

The negative-working image recording material of the present invention can perform recording independent of the emission wavelength of the exposing light source. In particular, the negative-working image recording material of the present invention can perform recording with light in the range of from near inferred to infrared (heat radiation).

The recording material of the present invention can perform recording by means of a solid laser or semiconductor laser (heat mode) having an emission wavelength range of from near infrared to infrared to make a plate directly from digital data from computer or the like. Further, a heat mode writing type direct plate making process with the recording material of the present invention which can make the direct application of conventional processors or printers can proside a lithorability printing the processors or printers can pro-

While the invention has been described in detail and with reference to specific embodiments thereof, it will be apparent to one skilled in the art that various changes and modifications can be made therein without departing from the scrift and accord thereof.

#### 10 Claims

A negative-working image recording material comprising:

(a) a substance which abscrbs light to generate heat:

(b) a water-insoluble and aqueous alkali-soluble resin; and

(c) a phenol derivative having in its molecule from 4 to 8 benzene nuclei, at least one phenolic hydroxyl group and at least two groups represented by the following general formula (f):

wherein R1 represents a hydrogen atom, an alkyl group or an acyl group.

- The negative-working image recording material as claimed in Claim 1, wherein said water-insoluble and aqueous alkali-soluble resin is a resin having a phenolic hydroxyl group or olefinically unsaturated bond.
- 3. A negative-working image recording material comprising a support having thereon (a) a substance which absorbs infrared rays to make infrared rays to generate heat, (b) a water-insoluble and aqueous salical positive teals in what penentic hydroxyl group or delificially unsaturated bond, and (a) a phenol derivative having in its molecule from 4 to 8 benzene nuclei, at least one phenolic hydroxyl group and at least two groups represented by the general formula (i):

wherein R1 represents a hydrogen atom, an alkyl group or an acyl group.

- The image recording material as claimed in Claim 3, wherein as said substance which absorbs infrared rays or near infrared rays to generate heat there is used a dive.
- The image recording material as claimed in Claim 3, wherein as said substance which absorbs infrared rays or near
   infrared rays to generate heat there is used a pigment.
  - The Image recording material as claimed in Claim 3, wherein as said substance which absorbs infrared rays or near infrared rays to generate heat there is used carbon black.
- 7. The image recording material as claimed in Claim 3, wherein as said water-insoluble and aqueous alkali-soluble resin having a phenolic hydroxyl group or olefinically unsaturated bond there is used a novolak resin.
  - The image recording material as dairmed in Claim 3, wherein as said water-insoluble and aqueous alkali-soluble resin having a phenolic hydroxyl group or olefinically unsaturated bond there is used an acrylic resin having an allyl group.
  - The image recording material as claimed in Claim 3, wherein as said water-insoluble and aqueous alkall-soluble rasin having a phenoic hydroxyl group or oldfinically unsaturated bond there is used an acrytic resin having a phenolic hydroxyl group.
  - The image recording material as claimed in Claim 3, wherein as said support there is used a polyester film.
    - 11. The image recording material as claimed in Claim 3, wherein as said support there is used an aluminum plate.

12. A negative-working image recording process, which comprises the steps of exposing a negative-working image recording meteral comprising a support having thereof, (a) a substance which absorbs infrared rays to generate heat. (b) a water-insoluble and aqueous allest escuble resid having a phenolic hydroxyl group or definically unsaturated bond, and (c) a phenol derivative having in its molecule from 4 to 6 benzene nuclei, at least one benefic hydroxyl group and at least the or pencilic hydroxyl group and a least which or groups represented by the general formula (f):

wherein R<sup>1</sup> represents a hydrogen atom, an alkyl group or an acyl group, to infrared or near infrared rays emitted by a laser, and then developing the negative-working image recording material with an aqueous alkali.

- The negative-working image recording process as claimed in Claim 12, wherein R<sup>1</sup> in the general formula (f) is an alkyl group.
- 14. The negative-working image recording process as claimed in Claim 12, wherein R<sup>1</sup> in the general formula (i) is a methyl group.
  - 15. The negative-working image recording process as claimed in Claim 12, further comprising the step of subjecting the negative-working image recording material to heat treatment between the exposing step and the developing step.

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#### (54) Negative-working image recording material

A negative-working image recording material is disclosed, which comprises a substance which absorbs light to generate heat, a water-insoluble and aqueous alkali-soluble resin, and a phenol derivative having in its molecule from 4 to 8 benzene nuclei, at least one phenotic hydroxyl group and at least two groups represented by the following general formula (I):

-CH<sub>2</sub>OR<sup>1</sup>

wherein R1 represents a hydrogen atom, an alkyl group or an acyl group. The negative-working image recording material can perform recording independent of the emission wavelength of the exposing light source, particularly with light in the range of from near infrared to infrared (heat radiation).



European Patent Office

# **EUROPEAN SEARCH REPORT**

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ategory	Citation of document with indicat of relevant passages	ion, where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.Cl.5)
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